

10/037, 519  
L3Cock 6/30/05

ANSWER 6 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
AN 1997:314054 BIOSIS  
DN PREV199799604542  
TI Stopped-flow kinetics reveal multiple phases of **thioflavin**  
**T** binding to Alzheimer beta(1-40) amyloid fibrils.  
AU Levine, Harry Iii  
CS Neurodegenerative Diseases, Parke-Davis Pharmaceutical Research Div.,  
Warner-Lambert Co., 2800 Plymouth Road, Ann Arbor, MI 48105-1047, USA  
SO Archives of Biochemistry and Biophysics, (1997) Vol. 342, No. 2, pp.  
306-316.  
CODEN: ABBIA4. ISSN: 0003-9861.  
DT Article  
LA English  
ED Entered STN: 26 Jul 1997  
Last Updated on STN: 4 Sep 1997  
AB The benzothiazole dye **thioflavin T** (ThT) is a  
classical amyloid stain for senile plaques containing beta/A4 peptide in  
Alzheimer's disease brain. ThT also binds rapidly and specifically to the  
anti-parallel beta-sheet fibrils formed from **synthetic**  
beta(1-40) peptide, but does not bind to monomer or oligomeric  
intermediates. The fibrillar beta-sheet-bound dye species undergoes a  
characteristic 120 nm red shift of its excitation spectrum that  
may be selectively excited at 450 nm, resulting in a  
fluorescence signal at 482 nm. Mixing of preformed beta(1-40)  
amyloid fibrils with ThT in a stopped-flow spectrophotometer, monitoring  
fluorescence emission at gt 475 nm while exciting at 450  
nm, distinguished multiple kinetic phases of roughly equivalent  
amplitude with tau's in the ranges of 0.007, 0.05, 0.75, and 10-20 s. The  
fastest reaction appears to reflect a bimolecular dye binding event while  
the remaining reactions are rate-limited by protein tertiary or quaternary  
conformational changes. The high activation energies of the three slower  
reactions support this interpretation. The ThT concentration dependence  
of the reaction rates at different ratios of ThT/beta(1-40) amyloid  
fibrils rules out a rate-limiting conformational change occurring prior to  
ligand binding. ThT is a useful probe for the **aggregated**  
fibrillar state of beta(1-40) amyloid fibrils as the amyloid-specific  
fluorescence reports only fibrillar species. The binding of ThT does not  
interfere with the **aggregation** of this peptide into amyloid  
fibrils. The putative conformational changes detected by the ThT  
fluorescence suggest that small pharmacologic ligands can perturb and  
possibly dissociate A-beta amyloid fibrils.  
CC Biochemistry studies - General 10060  
Biophysics - General 10502  
Nervous system - General and methods 20501  
IT Major Concepts  
Biochemistry and Molecular Biophysics; Nervous System (Neural  
Coordination)  
IT Chemicals & Biochemicals  
THIOFLAVIN; AMYLOID  
IT Time  
Quaternary; Tertiary  
IT Miscellaneous Descriptors  
ACTIVATION ENERGY; ALZHEIMER BETA(1-40) AMYLOID FIBRILS; ALZHEIMER'S  
DISEASE; BEHAVIORAL AND MENTAL DISORDERS; BIOCHEMISTRY AND BIOPHYSICS;  
BRAIN; CONFORMATIONAL CHANGES; KINETIC PHASES; NERVOUS SYSTEM; NERVOUS  
SYSTEM DISEASE; PHARMACOLOGIC LIGANDS; RATE-LIMITING; STOPPED-FLOW  
KINETICS; **THIOFLAVIN T** BINDING  
RN 2390-54-7 (THIOFLAVIN)  
11061-24-8 (AMYLOID)

ANSWER 9 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN

AN 1993:444464 CAPLUS

DN 119:44464

ED Entered STN: 07 Aug 1993

TI **Thioflavin T** interaction with **synthetic**  
Alzheimer's disease  $\beta$ -amyloid peptides: Detection of amyloid  
**aggregation** in solution

AU LeVine, Harry, III

CS Dep. Neurosci. Pharmacol., Warner-Lambert Co., Ann Arbor, MI, 48106-1047,  
USA

SO Protein Science (1993), 2(3), 404-10

CODEN: PRCIEI; ISSN: 0961-8368

DT Journal

LA English

CC 9-5 (Biochemical Methods)

Section cross-reference(s): 14

AB Thioflavine T (ThT) assoc. rapidly with **aggregated** fibrils of  
the **synthetic**  $\beta$ /A4-derived peptides  $\beta$ (1-28) and  
 $\beta$ (1-40), giving rise to a new excitation (ex) (absorption) maximum at  
450 nm and enhanced emission (em) at 482 nm, as  
opposed to the 385 nm (ex) and 445 nm (em) of the free  
dye. This change is dependent on the **aggregated** state as  
monomeric or dimeric peptides do not react, and guanidine dissociation of  
**aggregates** destroys the signal. There was no effect of high salt  
concns. Binding to the  $\beta$ (1-40) is of lower affinity,  $K_d$  2  $\mu$ M,  
while it sats. with a  $K_d$  of 0.54  $\mu$ M for  $\beta$ (1-28). Insulin fibrils  
converted to a  $\beta$ -sheet conformation fluoresce intensely with ThT. A  
variety of polyhydroxy, polyanionic, or polycationic materials fail to  
interact or impede interaction with the amyloid peptides. This  
fluorometric technique should allow the kinetic elucidation of the amyloid  
fibril assembly process as well as the testing of agents that might  
modulate their assembly or disassembly.

ST thioflavine T amyloid protein fluorescence Alzheimer

IT Mental disorder

(Alzheimer's disease, pathogenesis of, amyloid fibril formation in,  
thioflavine T interaction with **synthetic**  $\beta$ -amyloid  
protein-derived peptide fragments studied by fluorometry in relation  
to)

IT Proteins, specific or class

RL: ANST (Analytical study)

(amyloid A4, **synthetic** peptides derived from, thioflavine T  
interaction with, fluorometry in study of, Alzheimer's disease  
pathogenesis and amyloid fibril formation in relation to)

IT 2390-54-7, Thioflavine T

RL: ANST (Analytical study)

(**synthetic**  $\beta$ -amyloid peptide interaction with,  
fluorometry in study of, Alzheimer's disease pathogenesis and amyloid  
fibril formation in relation to)

10/037, 519  
L/COOK 6/30/05

=> s (thioflavin T)  
L1 1352 (THIOFLAVIN T)

=> s l1 and 485?  
TERM '485?' EXCEEDED TRUNCATION LIMITS - SEARCH ENDED  
You have entered a truncated stem which occurs in too many terms.  
Make the stem longer and try again. For example, if your original  
term was 'degr?' to search for variations and the abbreviation for  
'degradation', you could replace it with the expression '(degrdn OR  
degrad?)'. If your search term was numeric, e.g., 'C>5', reduce the  
size of the range.

=> s l1 and aggregat?  
L2 611 L1 AND AGGREGAT?

=> s l2 and nm?  
L3 121 L2 AND NM?

=> duplicate remove l3  
DUPLICATE PREFERENCE IS 'BIOSIS, CAPLUS, EMBASE, MEDLINE, CANCERLIT'  
KEEP DUPLICATES FROM MORE THAN ONE FILE? Y/(N):n  
PROCESSING COMPLETED FOR L3  
L4 45 DUPLICATE REMOVE L3 (76 DUPLICATES REMOVED)

=> s l4 and syn?  
L5 12 L4 AND SYN?

=> d l5 1-12 all

L5 ANSWER 1 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
AN 2003:513254 BIOSIS  
DN PREV200300516592  
TI Environmental influences on bovine kappa-casein: Reduction and conversion  
to fibrillar (amyloid) structures.  
AU Farrell, Harold M. Jr. [Reprint Author]; Cooke, Peter H.; Wickham, Edward  
D.; Piotrowski, Edwin G.; Hoagland, Peter D.  
CS Eastern Regional Research Center, United States Department of Agriculture,  
ARS, 600 E. Mermaid Lane, Wyndmoor, PA, 19038, USA  
hfarrell@arserrc.gov  
SO Journal of Protein Chemistry, (April 2003) Vol. 22, No. 3, pp. 259-273.  
print.  
ISSN: 0277-8033 (ISSN print).  
DT Article  
LA English  
ED Entered STN: 5 Nov 2003  
Last Updated on STN: 5 Nov 2003  
AB The caseins of milk form a unique calcium-phosphate transport complex that  
provides these necessary nutrients to the neonate. The colloidal  
stability of these particles is primarily the result of kappa-casein. As  
purified from milk, this protein occurs as spherical particles with a  
weight average molecular weight of 1.18 million. The protein exhibits a  
unique disulfide bonding pattern, which (in the absence of reducing  
agents) ranges from monomer to octamers and above on SDS-PAGE. Severe  
heat treatment of the kappa-casein (90degreeC) in the absence of SDS,  
before electrophoresis, caused an increase in the polymeric distribution:  
up to 40% randomly **aggregated** high-molecular weight polymers,  
presumably promoted by free sulfhydryl groups (J. Protein Chemical 17:  
73-84, 1998). To ascertain the role of the sulfhydryl groups, the protein  
was reduced and carboxymethylated (RCM-kappa). Surprisingly, at only  
37degreeC, the RCM-kappa-casein exhibited an increase in weight average  
molecular weight and tendency to self-association when studied at 3000 rpm  
by analytical ultracentrifugation. Electron microscopy (EM) of the  
37degreeC RCM sample showed that, in addition to the spherical particles

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L1 1352 (THIOFLAVIN T)

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Make the stem longer and try again. For example, if your original  
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degrad?)'. If your search term was numeric, e.g., 'C>5', reduce the  
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=> s l1 and aggregat?  
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=> s l2 and nm?  
L3 121 L2 AND NM?

=> duplicate remove l3  
DUPLICATE PREFERENCE IS 'BIOSIS, CAPLUS, EMBASE, MEDLINE, CANCERLIT'  
KEEP DUPLICATES FROM MORE THAN ONE FILE? Y/(N):n  
PROCESSING COMPLETED FOR L3  
L4 45 DUPLICATE REMOVE L3 (76 DUPLICATES REMOVED)

=> s l4 and syn?  
L5 12 L4 AND SYN?

=> d l5 1-12 all

L5 ANSWER 1 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
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TI Environmental influences on bovine kappa-casein: Reduction and conversion  
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AU Farrell, Harold M. Jr. [Reprint Author]; Cooke, Peter H.; Wickham, Edward  
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CS Eastern Regional Research Center, United States Department of Agriculture,  
ARS, 600 E. Mermaid Lane, Wyndmoor, PA, 19038, USA  
hfarrell@arserrc.gov  
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ISSN: 0277-8033 (ISSN print).  
DT Article  
LA English  
ED Entered STN: 5 Nov 2003  
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AB The caseins of milk form a unique calcium-phosphate transport complex that  
provides these necessary nutrients to the neonate. The colloidal  
stability of these particles is primarily the result of kappa-casein. As  
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weight average molecular weight of 1.18 million. The protein exhibits a  
unique disulfide bonding pattern, which (in the absence of reducing  
agents) ranges from monomer to octamers and above on SDS-PAGE. Severe  
heat treatment of the kappa-casein (90degreeC) in the absence of SDS,  
before electrophoresis, caused an increase in the polymeric distribution:  
up to 40% randomly **aggregated** high-molecular weight polymers,  
presumably promoted by free sulfhydryl groups (J. Protein Chemical 17:  
73-84, 1998). To ascertain the role of the sulfhydryl groups, the protein  
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37degreeC, the RCM-kappa-casein exhibited an increase in weight average  
molecular weight and tendency to self-association when studied at 3000 rpm  
by analytical ultracentrifugation. Electron microscopy (EM) of the  
37degreeC RCM sample showed that, in addition to the spherical particles

found in the native protein, there was a high proportion of fibrillar structures. The fibrillar structures were up to 600 nm in length. Circular dichroism (CD) spectroscopy was used to investigate the temperature-induced changes in the secondary structure of the native and RCM-kappa-caseins. These studies indicate that there was little change in the distribution of secondary structural elements during this transition, with extended strand and beta turns predominating. On the basis of three-dimensional molecular modeling predictions, there may exist a tyrosine-rich repeated sheet-turn-sheet motif in kappa-casein (residues 15-65), which may allow for the stacking of the molecules into fibrillar structures. Previous studies on amyloid proteins have suggested that such motifs promote fibril formation, and near-ultraviolet CD and **thioflavin-T** binding studies on RCM-kappa-casein support this concept. The results are discussed with respect to the role that such fibrils may play in the **synthesis** and secretion of casein micelles in lactating mammary gland.

CC Biochemistry studies - General 10060  
 Reproductive system - Physiology and biochemistry 16504  
 IT Major Concepts  
     Biochemistry and Molecular Biophysics; Reproductive System  
     (Reproduction)  
 IT Parts, Structures, & Systems of Organisms  
     mammary gland: reproductive system; milk: reproductive system  
 IT Chemicals & Biochemicals  
     kappa-casein: reduction, structure  
 IT Methods & Equipment  
     carboxymethylation: laboratory techniques; circular dichroism  
     spectroscopy: laboratory techniques, spectrum analysis techniques;  
     electron microscopy: imaging and microscopy techniques, laboratory  
     techniques; electrophoresis: electrophoretic techniques, laboratory  
     techniques; heat treatment: laboratory techniques; molecular modeling:  
     mathematical and computer techniques; reduction reaction: laboratory  
     techniques; ultracentrifugation: laboratory techniques  
 IT Miscellaneous Descriptors  
     kappa-casein fibril; lactation; temperature  
 ORGN Classifier  
     Bovidae 85715  
     Super Taxa  
     Artiodactyla; Mammalia; Vertebrata; Chordata; Animalia  
     Organism Name  
     bovine (common)  
     Taxa Notes  
     Animals, Artiodactyls, Chordates, Mammals, Nonhuman Vertebrates,  
     Nonhuman Mammals, Vertebrates

L5 ANSWER 2 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
 AN 2003:394261 BIOSIS  
 DN PREV200300394261  
 TI **Synthesis** and evaluation of 11C-labeled 6-substituted  
 2-arylbenzothiazoles as amyloid imaging agents.  
 AU Mathis, Chester A. [Reprint Author]; Wang, Yanming; Holt, Daniel P.;  
 Huang, Guo-feng; Debnath, Manik L.; Klunk, William E.  
 CS PET Facility, UPMC Presbyterian, 200 Lothrop Street, B-938, Pittsburgh,  
 PA, 15213-2582, USA  
 mathisca@msx.upmc.edu  
 SO Journal of Medicinal Chemistry, (June 19 2003) Vol. 46, No. 13, pp.  
 2740-2754. print.  
 ISSN: 0022-2623 (ISSN print).  
 DT Article  
 LA English  
 ED Entered STN: 27 Aug 2003  
 Last Updated on STN: 27 Aug 2003  
 AB The **synthesis** and evaluation of a series of neutral analogues of

found in the native protein, there was a high proportion of fibrillar structures. The fibrillar structures were up to 600 nm in length. Circular dichroism (CD) spectroscopy was used to investigate the temperature-induced changes in the secondary structure of the native and RCM-kappa-caseins. These studies indicate that there was little change in the distribution of secondary structural elements during this transition, with extended strand and beta turns predominating. On the basis of three-dimensional molecular modeling predictions, there may exist a tyrosine-rich repeated sheet-turn-sheet motif in kappa-casein (residues 15-65), which may allow for the stacking of the molecules into fibrillar structures. Previous studies on amyloid proteins have suggested that such motifs promote fibril formation, and near-ultraviolet CD and **thioflavin-T** binding studies on RCM-kappa-casein support this concept. The results are discussed with respect to the role that such fibrils may play in the **synthesis** and secretion of casein micelles in lactating mammary gland.

CC Biochemistry studies - General 10060  
 Reproductive system - Physiology and biochemistry 16504

IT Major Concepts  
 Biochemistry and Molecular Biophysics; Reproductive System  
 (Reproduction)

IT Parts, Structures, & Systems of Organisms  
 mammary gland: reproductive system; milk: reproductive system

IT Chemicals & Biochemicals  
 kappa-casein: reduction, structure

IT Methods & Equipment  
 carboxymethylation: laboratory techniques; circular dichroism  
 spectroscopy: laboratory techniques, spectrum analysis techniques;  
 electron microscopy: imaging and microscopy techniques, laboratory  
 techniques; electrophoresis: electrophoretic techniques, laboratory  
 techniques; heat treatment: laboratory techniques; molecular modeling:  
 mathematical and computer techniques; reduction reaction: laboratory  
 techniques; ultracentrifugation: laboratory techniques

IT Miscellaneous Descriptors  
 kappa-casein fibril; lactation; temperature

ORGN Classifier  
 Bovidae 85715  
 Super Taxa  
 Artiodactyla; Mammalia; Vertebrata; Chordata; Animalia  
 Organism Name  
 bovine (common)  
 Taxa Notes  
 Animals, Artiodactyls, Chordates, Mammals, Nonhuman Vertebrates,  
 Nonhuman Mammals, Vertebrates

L5 ANSWER 2 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
 AN 2003:394261 BIOSIS  
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 SO Journal of Medicinal Chemistry, (June 19 2003) Vol. 46, No. 13, pp.  
 2740-2754. print.  
 ISSN: 0022-2623 (ISSN print).  
 DT Article  
 LA English  
 ED Entered STN: 27 Aug 2003  
 Last Updated on STN: 27 Aug 2003  
 AB The **synthesis** and evaluation of a series of neutral analogues of

**thioflavin-T** (termed BTA's) with high affinities for **aggregated** amyloid and a wide range of lipophilicities are reported. Radiolabeling with high specific activity (<sup>11</sup>C)methyl iodide provided derivatives for in vivo evaluation. Brain entry in control mice and baboons was high for nearly all of the analogues at early times after injection, but the clearance rate of radioactivity from brain tissue varied by more than 1 order of magnitude. Upon the basis of its rapid clearance from normal mouse and baboon brain tissues, (N-methyl-<sup>11</sup>C)2-(4'-methylaminophenyl)-6-hydroxybenzothiazole (or (<sup>11</sup>C)6-OH-BTA-1) was selected as the lead compound for further evaluation. The radiolabeled metabolites of (<sup>11</sup>C)6-OH-BTA-1 were polar and did not enter brain. The binding affinities of (N-methyl-<sup>3</sup>H)6-OH-BTA-1 for homogenates of postmortem AD frontal cortex and **synthetic** Abeta(1-40) fibrils were similar (K<sub>d</sub>=1.4 nM and 4.7 nM, respectively), but the ligand-to-Abeta peptide binding stoichiometry was approx 400-fold higher for AD brain than Abeta(1-40) fibrils. Staining of AD frontal cortex tissue sections with 6-OH-BTA-1 indicated the selective binding of the compound to amyloid plaques and cerebrovascular amyloid. The encouraging in vitro and in vivo properties of (<sup>11</sup>C)6-OH-BTA-1 support the choice of this derivative for further evaluation in human subject studies of brain Abeta deposition.

CC Behavioral biology - Human behavior 07004  
 Pathology - Diagnostic 12504  
 Pathology - Therapy 12512  
 Nervous system - Physiology and biochemistry 20504  
 Nervous system - Pathology 20506  
 Psychiatry - Psychopathology, psychodynamics and therapy 21002  
 Pharmacology - General 22002  
 Pharmacology - Clinical pharmacology 22005

IT Major Concepts  
 Methods and Techniques; Nervous System (Neural Coordination);  
 Pharmacology

IT Parts, Structures, & Systems of Organisms  
 brain: nervous system; frontal cortex: nervous system

IT Diseases  
 Alzheimer's disease: behavioral and mental disorders, nervous system  
 disease  
 Alzheimer Disease (MeSH)

IT Chemicals & Biochemicals  
 amyloid-beta; carbon-11-labeled 6-substituted 2-arylbenzothiazole:  
 diagnostic-drug, imaging agent, **synthesis**

IT Methods & Equipment  
 radiolabeling: laboratory techniques

IT Miscellaneous Descriptors  
 amyloid plaques; amyloid-beta(1-40) fibril; cerebrovascular amyloid

ORGN Classifier  
 Cercopithecidae 86205  
 Super Taxa  
 Primates; Mammalia; Vertebrata; Chordata; Animalia  
 Organism Name  
 Papio anubis (species) [baboon (common)]  
 Taxa Notes  
 Animals, Chordates, Mammals, Nonhuman Mammals, Nonhuman Vertebrates,  
 Nonhuman Primates, Primates, Vertebrates

ORGN Classifier  
 Hominidae 86215  
 Super Taxa  
 Primates; Mammalia; Vertebrata; Chordata; Animalia  
 Organism Name  
 human (common)  
 Taxa Notes  
 Animals, Chordates, Humans, Mammals, Primates, Vertebrates

ORGN Classifier

**thioflavin-T** (termed BTA's) with high affinities for **aggregated** amyloid and a wide range of lipophilicities are reported. Radiolabeling with high specific activity (11C)methyl iodide provided derivatives for in vivo evaluation. Brain entry in control mice and baboons was high for nearly all of the analogues at early times after injection, but the clearance rate of radioactivity from brain tissue varied by more than 1 order of magnitude. Upon the basis of its rapid clearance from normal mouse and baboon brain tissues, (N-methyl-11C)2-(4'-methylaminophenyl)-6-hydroxybenzothiazole (or (11C)6-OH-BTA-1) was selected as the lead compound for further evaluation. The radiolabeled metabolites of (11C)6-OH-BTA-1 were polar and did not enter brain. The binding affinities of (N-methyl-3H)6-OH-BTA-1 for homogenates of postmortem AD frontal cortex and **synthetic** Abeta(1-40) fibrils were similar ( $K_d=1.4$  nM and 4.7 nM, respectively), but the ligand-to-Abeta peptide binding stoichiometry was approx400-fold higher for AD brain than Abeta(1-40) fibrils. Staining of AD frontal cortex tissue sections with 6-OH-BTA-1 indicated the selective binding of the compound to amyloid plaques and cerebrovascular amyloid. The encouraging in vitro and in vivo properties of (11C)6-OH-BTA-1 support the choice of this derivative for further evaluation in human subject studies of brain Abeta deposition.

CC Behavioral biology - Human behavior 07004  
 Pathology - Diagnostic 12504  
 Pathology - Therapy 12512  
 Nervous system - Physiology and biochemistry 20504  
 Nervous system - Pathology 20506  
 Psychiatry - Psychopathology; psychodynamics and therapy 21002  
 Pharmacology - General 22002  
 Pharmacology - Clinical pharmacology 22005

IT Major Concepts  
 Methods and Techniques; Nervous System (Neural Coordination);  
 Pharmacology

IT Parts, Structures, & Systems of Organisms  
 brain: nervous system; frontal cortex: nervous system

IT Diseases  
 Alzheimer's disease: behavioral and mental disorders, nervous system  
 disease  
 Alzheimer Disease (MeSH)

IT Chemicals & Biochemicals  
 amyloid-beta; carbon-11-labeled 6-substituted 2-arylbenzothiazole:  
 diagnostic-drug, imaging agent, **synthesis**

IT Methods & Equipment  
 radiolabeling: laboratory techniques

IT Miscellaneous Descriptors  
 amyloid plaques; amyloid-beta(1-40) fibril; cerebrovascular amyloid

ORGN Classifier  
 Cercopithecidae 86205  
 Super Taxa  
 Primates; Mammalia; Vertebrata; Chordata; Animalia  
 Organism Name  
 Papio anubis (species) [baboon (common)]  
 Taxa Notes  
 Animals, Chordates, Mammals, Nonhuman Mammals, Nonhuman Vertebrates,  
 Nonhuman Primates, Primates, Vertebrates

ORGN Classifier  
 Hominidae 86215  
 Super Taxa  
 Primates; Mammalia; Vertebrata; Chordata; Animalia  
 Organism Name  
 human (common)  
 Taxa Notes  
 Animals, Chordates, Humans, Mammals, Primates, Vertebrates

ORGN Classifier



Muridae 86375  
Super Taxa  
Rodentia; Mammalia; Vertebrata; Chordata; Animalia  
Organism Name  
mouse (common)  
Taxa Notes  
Animals, Chordates, Mammals, Nonhuman Vertebrates, Nonhuman Mammals,  
Rodents, Vertebrates

L5 ANSWER 3 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
AN 2003:267944 BIOSIS  
DN PREV200300267944  
TI METAL - DEPENDENCE OF A beta OLIGOMERIZATION.  
AU Huang, X. [Reprint Author]; Moir, R. D.; Friedlich, A. L. [Reprint  
Author]; Nagano, S. [Reprint Author]; Goldstein, L. E. [Reprint Author];  
Rogers, J. T. [Reprint Author]; Tanzi, R. E.; Bush, A. I. [Reprint Author]  
CS Psychiatry/Genetics and Aging Research Unit, MGH/ Harvard Medical School,  
Charlestown, MA, USA  
SO Society for Neuroscience Abstract Viewer and Itinerary Planner, (2002)  
Vol. 2002, pp. Abstract No. 19.1. <http://sfn.scholarone.com>. cd-rom.  
Meeting Info.: 32nd Annual Meeting of the Society for Neuroscience.  
Orlando, Florida, USA. November 02-07, 2002. Society for Neuroscience.  
DT Conference; (Meeting)  
Conference; (Meeting Poster)  
Conference; Abstract; (Meeting Abstract)  
LA English  
ED Entered STN: 11 Jun 2003  
Last Updated on STN: 11 Jun 2003  
AB Introduction: Recent studies have demonstrated that diffusible human A  
oligomers, i.e. Abeta-derived diffusible ligands (ADDLs) are neurotoxic.  
It is known that iron, copper, and zinc are highly enriched in amyloid  
plaques. We have previously found that these metal ions are involved in  
maintaining the assembly of Abeta amyloid in vitro, and in post-mortem  
Alzheimer Disease (AD) brain specimens. We recently discovered that  
treatment with BBB-permeable metal chelator-clioquinol (CQ) inhibited  
Abeta deposition in APP2576 transgenic mice. Here we study the effects of  
these metal ions and CQ upon the Abeta oligomerization process used to  
form ADDLs. Methods: Metal concentrations in cold F12 medium were  
determined by ICP-MS. Abeta40 and Abeta42 (10 muM) in cold F12 medium  
were co-incubated at 4C 5 muM CQ or DTPA (another potent chelator).  
Turbidity readings (400 nm) were taken daily over ten days. At  
the time point where turbidity values plateaued, Abeta **aggregation**  
was quantified by Congo-Red and **Thioflavin-T** assays.  
The ADDLs were appraised by protein gel staining and FPLC. Results and  
Conclusion: The medium was found to contain 0.3 muM of copper, 16 muM of  
zinc, and 34 muM of iron. We observed the attenuation of Abeta  
oligomerization by both CQ and DTPA. Hence, the formation of ADDLs is  
induced by the presence of these metal ions in the medium. Specific metal  
chelators may be therapeutic for AD in interdicting **synaptotoxic**  
Abeta oligomerization.  
CC General biology - Symposia, transactions and proceedings 00520  
Biochemistry studies - Minerals 10069  
Pathology - Therapy 12512  
Nervous system - Physiology and biochemistry 20504  
Nervous system - Pathology 20506  
IT Major Concepts  
Nervous System (Neural Coordination)  
IT Diseases  
Alzheimer disease: behavioral and mental disorders, nervous system  
disease, therapy  
Alzheimer Disease (MeSH)  
IT Chemicals & Biochemicals  
A-beta [amyloid-beta]: **synaptotoxic**, deposition,

Muridae 86375  
Super Taxa  
Rodentia; Mammalia; Vertebrata; Chordata; Animalia  
Organism Name  
mouse (common)  
Taxa Notes  
Animals, Chordates, Mammals, Nonhuman Vertebrates, Nonhuman Mammals,  
Rodents, Vertebrates

L5 ANSWER 3 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
AN 2003:267944 BIOSIS  
DN PREV200300267944  
TI METAL - DEPENDENCE OF A beta OLIGOMERIZATION.  
AU Huang, X. [Reprint Author]; Moir, R. D.; Friedlich, A. L. [Reprint  
Author]; Nagano, S. [Reprint Author]; Goldstein, L. E. [Reprint Author];  
Rogers, J. T. [Reprint Author]; Tanzi, R. E.; Bush, A. I. [Reprint Author]  
CS Psychiatry/Genetics and Aging Research Unit, MGH/ Harvard Medical School,  
Charlestown, MA, USA  
SO Society for Neuroscience Abstract Viewer and Itinerary Planner, (2002)  
Vol. 2002, pp. Abstract No. 19.1. <http://sfn.scholarone.com>. cd-rom.  
Meeting Info.: 32nd Annual Meeting of the Society for Neuroscience.  
Orlando, Florida, USA. November 02-07, 2002. Society for Neuroscience.  
DT Conference; (Meeting)  
Conference; (Meeting Poster)  
Conference; Abstract; (Meeting Abstract)  
LA English  
ED Entered STN: 11 Jun 2003  
Last Updated on STN: 11 Jun 2003  
AB Introduction: Recent studies have demonstrated that diffusible human A  
oligomers, i.e. Abeta-derived diffusible ligands (ADDLs) are neurotoxic.  
It is known that iron, copper, and zinc are highly enriched in amyloid  
plaques. We have previously found that these metal ions are involved in  
maintaining the assembly of Abeta amyloid in vitro, and in post-mortem  
Alzheimer Disease (AD) brain specimens. We recently discovered that  
treatment with BBB-permeable metal chelator-clioquinol (CQ) inhibited  
Abeta deposition in APP2576 transgenic mice. Here we study the effects of  
these metal ions and CQ upon the Abeta oligomerization process used to  
form ADDLs. Methods: Metal concentrations in cold F12 medium were  
determined by ICP-MS. Abeta40 and Abeta42 (10 muM) in cold F12 medium  
were co-incubated at 4C 5 muM CQ or DTPA (another potent chelator).  
Turbidity readings (400 nm) were taken daily over ten days. At  
the time point where turbidity values plateaued, Abeta **aggregation**  
was quantified by Congo-Red and **Thioflavin-T** assays.  
The ADDLs were appraised by protein gel staining and FPLC. Results and  
Conclusion: The medium was found to contain 0.3 muM of copper, 16 muM of  
zinc, and 34 muM of iron. We observed the attenuation of Abeta  
oligomerization by both CQ and DTPA. Hence, the formation of ADDLs is  
induced by the presence of these metal ions in the medium. Specific metal  
chelators may be therapeutic for AD in interdicting **synaptotoxic**  
Abeta oligomerization.  
CC General biology - Symposia, transactions and proceedings 00520  
Biochemistry studies - Minerals 10069  
Pathology - Therapy 12512  
Nervous system - Physiology and biochemistry 20504  
Nervous system - Pathology 20506  
IT Major Concepts  
Nervous System (Neural Coordination)  
IT Diseases  
Alzheimer disease: behavioral and mental disorders, nervous system  
disease, therapy  
Alzheimer Disease (MeSH)  
IT Chemicals & Biochemicals  
A-beta [amyloid-beta]: **synaptotoxic**, deposition,

oligomerization; A-beta 40 [amyloid-beta 40]; A-beta 42 [amyloid-beta 42]; DTPA; F12: medium; amyloid beta-derived diffusible ligand [ADDL]; clioquinol: chelating agent; copper; iron; zinc

IT Miscellaneous Descriptors  
turbidity reading

ORGN Classifier  
Muridae 86375  
Super Taxa  
Rodentia; Mammalia; Vertebrata; Chordata; Animalia  
Organism Name  
mouse (common)  
Taxa Notes  
Animals, Chordates, Mammals, Nonhuman Vertebrates, Nonhuman Mammals, Rodents, Vertebrates

RN 67-43-6 (DTPA)  
130-26-7 (clioquinol)  
7440-50-8 (copper)  
7439-89-6 (iron)  
7440-66-6 (zinc)

L5 ANSWER 4 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
AN 2003:48300 BIOSIS  
DN PREV200300048300  
TI IMPY: An improved **thioflavin-T** derivative for in vivo labeling of beta-amyloid plaques.  
AU Kung, Mei-Ping [Reprint Author]; Hou, Catherine; Zhuang, Zhi-Ping; Zhang, Bin; Skovronsky, Daniel; Trojanowski, John Q.; Lee, Virginia M.-Y.; Kung, Hank F.  
CS Department of Radiology, University of Pennsylvania, 3700 Market Street, Room 305, Philadelphia, PA, 19104, USA  
kungmp@sunmac.spect.upenn.edu  
SO Brain Research, (29 November 2002) Vol. 956, No. 2, pp. 202-210. print. ISSN: 0006-8993 (ISSN print).  
DT Article  
LA English  
ED Entered STN: 15 Jan 2003  
Last Updated on STN: 15 Jan 2003  
AB Development of small molecular probes for in vivo labeling and detection of beta-amyloid (Abeta) plaques in patients of Alzheimer's disease (AD) is of significant scientific interest, and it may also assist the development of drugs targeting Abeta plaques for treatment of AD. A novel probe, (123I/125I)IMPY, 6-iodo-2-(4'-dimethylamino-)phenyl-imidazo(1,2-a)pyridine, was successfully prepared with an iododestannylation reaction catalyzed by hydrogen peroxide. The modified **thioflavin-T** derivative displayed a good binding affinity for preformed **synthetic Abeta40 aggregates** in solution ( $K_i=15+-5$  nM) and showed selective plaque labeling on postmortem AD brain sections. Biodistribution study in normal mice after an iv injection of (125I)IMPY exhibited excellent brain uptake (2.9% initial dose/brain at 2 min) and fast washout (0.2% initial dose/brain at 60 min). These properties are highly desirable for amyloid plaque imaging agents. In vivo plaque labeling was evaluated in a transgenic mouse model (Tg2576) engineered to produce excess amyloid plaques in the brain. Ex vivo autoradiograms of brain sections of the Tg 2576 mouse obtained at 4 h after an i.v. injection of (125I)IMPY clearly displayed a distinct plaque labeling with a low background activity. When the same brain section was stained with a fluorescent dye, thioflavin-S, the same Abeta plaques showed prominent fluorescent labeling consistent with the results of the autoradiogram. In conclusion, these findings clearly suggest that radioiodinated IMPY demonstrates desirable characteristics for in vivo labeling of Abeta plaques and it may be useful as a molecular imaging agent to study amyloidogenesis in the brain of living AD patients.

CC Behavioral biology - Human behavior 07004

oligomerization; A-beta 40 [amyloid-beta 40]; A-beta 42 [amyloid-beta 42]; DTPA; F12: medium; amyloid beta-derived diffusible ligand [ADDL]; clioquinol: chelating agent; copper; iron; zinc

IT Miscellaneous Descriptors  
turbidity reading

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Muridae 86375  
Super Taxa  
Rodentia; Mammalia; Vertebrata; Chordata; Animalia  
Organism Name  
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Taxa Notes  
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7440-50-8 (copper)  
7439-89-6 (iron)  
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CC Behavioral biology - Human behavior 07004

Biochemistry studies - General 10060  
 Biochemistry studies - Proteins, peptides and amino acids 10064  
 Nervous system - Physiology and biochemistry 20504  
 Nervous system - Pathology 20506  
 Psychiatry - Psychopathology, psychodynamics and therapy 21002  
 IT Major Concepts  
     Nervous System (Neural Coordination)  
 IT Parts, Structures, & Systems of Organisms  
     beta-amyloid plaque: nervous system; brain: nervous system  
 IT Diseases  
     Alzheimer's disease: behavioral and mental disorders, nervous system disease  
     Alzheimer Disease (MeSH)  
 IT Chemicals & Biochemicals  
     IMPY: molecular probe; beta-amyloid; hydrogen peroxide;  
     **thioflavin-T**  
 IT Methods & Equipment  
     in vivo labeling: laboratory techniques  
 IT Miscellaneous Descriptors  
     amyloidogenesis  
 ORGN Classifier  
     Hominidae 86215  
     Super Taxa  
         Primates; Mammalia; Vertebrata; Chordata; Animalia  
     Organism Name  
         human (common): patient  
     Taxa Notes  
         Animals, Chordates, Humans, Mammals, Primates, Vertebrates  
 ORGN Classifier  
     Muridae 86375  
     Super Taxa  
         Rodentia; Mammalia; Vertebrata; Chordata; Animalia  
     Organism Name  
         mouse (common): animal model, transgenic  
     Taxa Notes  
         Animals, Chordates, Mammals, Nonhuman Vertebrates, Nonhuman Mammals, Rodents, Vertebrates  
 RN 7722-84-1 (hydrogen peroxide)  
     2390-54-7 (**thioflavin-T**)  
  
 L5 ANSWER 5 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
 AN 2003:43126 BIOSIS  
 DN PREV200300043126  
 TI Amyloid fibril formation by a **synthetic** peptide from a region of human acetylcholinesterase that is homologous to the Alzheimer's amyloid-beta peptide.  
 AU Cottingham, Matthew G.; Hollinshead, Michael S.; Vaux, David J. T.  
     [Reprint Author]  
 CS Sir William Dunn School of Pathology, University of Oxford, Oxford, OX1 3RE, UK  
     vaux@molbiol.ox.ac.uk  
 SO Biochemistry, (November 19 2002) Vol. 41, No. 46, pp. 13539-13547. print.  
     ISSN: 0006-2960 (ISSN print).  
 DT Article  
 LA English  
 ED Entered STN: 15 Jan 2003  
     Last Updated on STN: 15 Jan 2003  
 AB A region near the C-terminus of human acetylcholinesterase (AChE) is weakly homologous with the N-terminus of the Alzheimer's disease amyloid-beta peptide. We report that a 14-amino acid **synthetic** polypeptide whose sequence corresponds to residues 586-599 of the human **synaptic** or T form of AChE assembles into amyloid fibrils under physiological conditions. The fibrils have all the classical

Biochemistry studies - General 10060  
 Biochemistry studies - Proteins, peptides and amino acids 10064  
 Nervous system - Physiology and biochemistry 20504  
 Nervous system - Pathology 20506  
 Psychiatry - Psychopathology, psychodynamics and therapy 21002  
 IT Major Concepts  
     Nervous System (Neural Coordination)  
 IT Parts, Structures, & Systems of Organisms  
     beta-amyloid plaque: nervous system; brain: nervous system  
 IT Diseases  
     Alzheimer's disease: behavioral and mental disorders, nervous system disease  
     Alzheimer Disease (MeSH)  
 IT Chemicals & Biochemicals  
     IMPY: molecular probe; beta-amyloid; hydrogen peroxide;  
     **thioflavin-T**  
 IT Methods & Equipment  
     in vivo labeling: laboratory techniques  
 IT Miscellaneous Descriptors  
     amyloidogenesis  
 ORGN Classifier  
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     Super Taxa  
         Primates; Mammalia; Vertebrata; Chordata; Animalia  
     Organism Name  
         human (common): patient  
     Taxa Notes  
         Animals, Chordates, Humans, Mammals, Primates, Vertebrates  
 ORGN Classifier  
     Muridae 86375  
     Super Taxa  
         Rodentia; Mammalia; Vertebrata; Chordata; Animalia  
     Organism Name  
         mouse (common): animal model, transgenic  
     Taxa Notes  
         Animals, Chordates, Mammals, Nonhuman Vertebrates, Nonhuman Mammals, Rodents, Vertebrates  
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     2390-54-7 (**thioflavin-T**)  
  
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characteristics of amyloid: they have a diameter of 6-7 nm and bind both Congo red and **thioflavin-T**. Furthermore, the kinetics of assembly indicate that fibril formation proceeds via a two-step nucleation-dependent polymerization pathway, and a transition in the peptide conformation from random coil to beta-sheet is observed during fibril formation using far-UV circular dichroism spectroscopy. We also show that the peptide in **aggregated** fibrillar form has a toxic effect upon PC-12 cells in vitro. AChE normally resides mainly on cholinergic neuronal membranes, but is abnormally localized to senile plaques in Alzheimer's disease. Recently, an in vitro interaction between AChE and Abeta, the principal constituent of the amyloid fibrils in senile plaques, has been documented. The presence of a fibrillogenic region within AChE may be relevant to the interaction of AChE with amyloid fibrils formed by Abeta.

CC Behavioral biology - Human behavior 07004  
 Biochemistry studies - General 10060  
 Biochemistry studies - Proteins, peptides and amino acids 10064  
 Enzymes - General and comparative studies: coenzymes 10802  
 Nervous system - Physiology and biochemistry 20504  
 Nervous system - Pathology 20506  
 Psychiatry - Psychopathology, psychodynamics and therapy 21002  
 Toxicology - General and methods 22501

IT Major Concepts  
 Biochemistry and Molecular Biophysics; Nervous System (Neural Coordination)

IT Diseases  
 Alzheimer's disease: behavioral and mental disorders, nervous system disease  
 Alzheimer Disease (MeSH)

IT Chemicals & Biochemicals  
 acetylcholinesterase [EC 3.1.1.7]: activities, functions, human, molecular analysis; amyloid fibrils: analysis, formation; amyloid-beta peptide; enzymes; peptides; proteins; **synthetic** peptides

IT Methods & Equipment  
 far-UV circular dichroism spectroscopy: laboratory techniques, spectrum analysis techniques

IT Miscellaneous Descriptors  
 comparative biochemistry; molecular interactions; neuropathology; physiological conditions; toxicity

ORGN Classifier  
 Hominidae 86215  
 Super Taxa  
 Primates; Mammalia; Vertebrata; Chordata; Animalia  
 Organism Name  
 human (common)  
 Taxa Notes  
 Animals, Chordates, Humans, Mammals, Primates, Vertebrates

ORGN Classifier  
 Muridae 86375  
 Super Taxa  
 Rodentia; Mammalia; Vertebrata; Chordata; Animalia  
 Organism Name  
 PC-12 cell line (cell line)  
 Taxa Notes  
 Animals, Chordates, Mammals, Nonhuman Vertebrates, Nonhuman Mammals, Rodents, Vertebrates

RN 9000-81-1 (acetylcholinesterase)  
 9000-81-1 (EC 3.1.1.7)

L5 ANSWER 6 OF 12 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN  
 AN 1997:314054 BIOSIS  
 DN PREV199799604542  
 TI Stopped-flow kinetics reveal multiple phases of **thioflavin**

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CC Behavioral biology - Human behavior 07004  
 Biochemistry studies - General 10060  
 Biochemistry studies - Proteins, peptides and amino acids 10064  
 Enzymes - General and comparative studies: coenzymes 10802  
 Nervous system - Physiology and biochemistry 20504  
 Nervous system - Pathology 20506  
 Psychiatry - Psychopathology, psychodynamics and therapy 21002  
 Toxicology - General and methods 22501

IT Major Concepts  
 Biochemistry and Molecular Biophysics; Nervous System (Neural Coordination)

IT Diseases  
 Alzheimer's disease: behavioral and mental disorders, nervous system disease  
 Alzheimer Disease (MeSH)

IT Chemicals & Biochemicals  
 acetylcholinesterase [EC 3.1.1.7]: activities, functions, human, molecular analysis; amyloid fibrils: analysis, formation; amyloid-beta peptide; enzymes; peptides; proteins; **synthetic** peptides

IT Methods & Equipment  
 far-UV circular dichroism spectroscopy: laboratory techniques, spectrum analysis techniques

IT Miscellaneous Descriptors  
 comparative biochemistry; molecular interactions; neuropathology; physiological conditions; toxicity

ORGN Classifier  
 Hominidae 86215  
 Super Taxa  
 Primates; Mammalia; Vertebrata; Chordata; Animalia  
 Organism Name  
 human (common)  
 Taxa Notes  
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 AN 1997:314054 BIOSIS  
 DN PREV199799604542  
 TI Stopped-flow kinetics reveal multiple phases of **thioflavin**



T binding to Alzheimer beta(1-40) amyloid fibrils.

AU Levine, Harry Iii

CS Neurodegenerative Diseases, Parke-Davis Pharmaceutical Research Div.,  
Warner-Lambert Co., 2800 Plymouth Road, Ann Arbor, MI 48105-1047, USA

SO Archives of Biochemistry and Biophysics, (1997) Vol. 342, No. 2, pp.  
306-316.  
CODEN: ABBIA4. ISSN: 0003-9861.

DT Article

LA English

ED Entered STN: 26 Jul 1997  
Last Updated on STN: 4 Sep 1997

AB The benzothiazole dye **thioflavin T** (ThT) is a classical amyloid stain for senile plaques containing beta/A4 peptide in Alzheimer's disease brain. ThT also binds rapidly and specifically to the anti-parallel beta-sheet fibrils formed from **synthetic** beta(1-40) peptide, but does not bind to monomer or oligomeric intermediates. The fibrillar beta-sheet-bound dye species undergoes a characteristic 120 nm red shift of its excitation spectrum that may be selectively excited at 450 nm, resulting in a fluorescence signal at 482 nm. Mixing of preformed beta(1-40) amyloid fibrils with ThT in a stopped-flow spectrophotometer, monitoring fluorescence emission at gt 475 nm while exciting at 450 nm, distinguished multiple kinetic phases of roughly equivalent amplitude with tau's in the ranges of 0.007, 0.05, 0.75, and 10-20 s. The fastest reaction appears to reflect a bimolecular dye binding event while the remaining reactions are rate-limited by protein tertiary or quaternary conformational changes. The high activation energies of the three slower reactions support this interpretation. The ThT concentration dependence of the reaction rates at different ratios of ThT/beta(1-40) amyloid fibrils rules out a rate-limiting conformational change occurring prior to ligand binding. ThT is a useful probe for the **aggregated** fibrillar state of beta(1-40) amyloid fibrils as the amyloid-specific fluorescence reports only fibrillar species. The binding of ThT does not interfere with the **aggregation** of this peptide into amyloid fibrils. The putative conformational changes detected by the ThT fluorescence suggest that small pharmacologic ligands can perturb and possibly dissociate A-beta amyloid fibrils.

CC Biochemistry studies - General 10060  
Biophysics - General 10502  
Nervous system - General and methods 20501

IT Major Concepts  
Biochemistry and Molecular Biophysics; Nervous System (Neural Coordination)

IT Chemicals & Biochemicals  
THIOFLAVIN; AMYLOID

IT Time  
Quaternary; Tertiary

IT Miscellaneous Descriptors  
ACTIVATION ENERGY; ALZHEIMER BETA(1-40) AMYLOID FIBRILS; ALZHEIMER'S DISEASE; BEHAVIORAL AND MENTAL DISORDERS; BIOCHEMISTRY AND BIOPHYSICS; BRAIN; CONFORMATIONAL CHANGES; KINETIC PHASES; NERVOUS SYSTEM; NERVOUS SYSTEM DISEASE; PHARMACOLOGIC LIGANDS; RATE-LIMITING; STOPPED-FLOW KINETICS; **THIOFLAVIN T BINDING**

RN 2390-54-7 (THIOFLAVIN)  
11061-24-8 (AMYLOID)

L5 ANSWER 7 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN

AN 2005:327565 CAPLUS

ED Entered STN: 18 Apr 2005

TI Neurotoxic effect of rotenone on dopaminergic neurons

AU Qi, Chen; Liu, Zhenguo; Fan, Guohua; Chen, Shengdi; Lu, Guoqiang

CS Ruijin Hospital, Shanghai Second Medical University, Shanghai, 200025, Peop. Rep. China

T binding to Alzheimer beta(1-40) amyloid fibrils.

AU Levine, Harry Iii  
CS Neurodegenerative Diseases, Parke-Davis Pharmaceutical Research Div.,  
Warner-Lambert Co., 2800 Plymouth Road, Ann Arbor, MI 48105-1047, USA  
SO Archives of Biochemistry and Biophysics, (1997) Vol. 342, No. 2, pp.  
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fluorescence reports only fibrillar species. The binding of ThT does not  
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fibrils. The putative conformational changes detected by the ThT  
fluorescence suggest that small pharmacologic ligands can perturb and  
possibly dissociate A-beta amyloid fibrils.  
CC Biochemistry studies - General 10060  
Biophysics - General 10502  
Nervous system - General and methods 20501  
IT Major Concepts  
Biochemistry and Molecular Biophysics; Nervous System (Neural  
Coordination)  
IT Chemicals & Biochemicals  
THIOFLAVIN; AMYLOID  
IT Time  
Quaternary; Tertiary  
IT Miscellaneous Descriptors  
ACTIVATION ENERGY; ALZHEIMER BETA(1-40) AMYLOID FIBRILS; ALZHEIMER'S  
DISEASE; BEHAVIORAL AND MENTAL DISORDERS; BIOCHEMISTRY AND BIOPHYSICS;  
BRAIN; CONFORMATIONAL CHANGES; KINETIC PHASES; NERVOUS SYSTEM; NERVOUS  
SYSTEM DISEASE; PHARMACOLOGIC LIGANDS; RATE-LIMITING; STOPPED-FLOW  
KINETICS; **THIOFLAVIN T BINDING**  
RN 2390-54-7 (THIOFLAVIN)  
11061-24-8 (AMYLOID)  
L5 ANSWER 7 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN  
AN 2005:327565 CAPLUS  
ED Entered STN: 18 Apr 2005  
TI Neurotoxic effect of rotenone on dopaminergic neurons  
AU Qi, Chen; Liu, Zhenguo; Fan, Guohua; Chen, Shengdi; Lu, Guoqiang  
CS Ruijin Hospital, Shanghai Second Medical University, Shanghai, 200025,  
Peop. Rep. China

SO Zhonghua Shenjingke Zazhi (2004), 37(6), 538-542  
 CODEN: ZSZAFN; ISSN: 1006-7876  
 PB Zhonghua Yixuehui Zazhishe  
 DT Journal  
 LA Chinese  
 CC 4 (Toxicology)  
 AB The mechanism of rotenone neurotoxicity on dopaminergic neurons was investigated. PC12 cells differentiated by nerve growth factor as dopaminergic neurons were treated by different concns. of rotenone. Cell viability was assessed with MTT, and cell apoptosis was detected by Annexin-V staining and flow cytometry. The double staining with  $\alpha$ -**synuclein** and **thioflavin T** was used to observe protein **aggregation**. After being treated with rotenone for 24 h, the process-like structures of PC12 cells disappeared, and the cell body became smaller and smoother in time- and concentration-dependent manners. Compared with the control group, the cell viability began to decline significantly when treated by rotenone at concentration of 10 **nmol/L** ( $A_{570} 0.415 \pm 0.013$ ) ( $P < 0.05$ ). The early sign of apoptosis was found with Annexin-V pos. staining. The apoptotic rate was  $7.35 \pm 0.52\%$  at rotenone concentration of 5 **nmol/L** ( $P < 0.05$ ), and was  $13.30 \pm 1.80\%$  at concentration of 10 **nmol/L** ( $P < 0.01$ ). Protein **aggregation** with the double pos. staining of  $\alpha$ - **synuclein** and **thioflavin T** were also found in the groups treated by rotenone. In vitro, rotenone should be neurotoxic to dopaminergic neurons, inducing apoptosis and inclusion of  $\alpha$ - **synuclein aggregation**. Rotenone might act through the metabolism of  $\alpha$ - **synuclein** in the pathogenesis of Parkinson's disease.  
 ST rotenone neurotoxicity dopaminergic neuron Parkinson disease

L5 ANSWER 8 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN  
 AN 2001:827673 CAPLUS  
 DN 137:59572  
 ED Entered STN: 14 Nov 2001  
 TI IBOX (2-(4'-dimethylaminophenyl)-6-iodobenzoxazole): a ligand for imaging amyloid plaques in the brain  
 AU Zhuang, Zhi-Ping; Kung, Mei-Ping; Hou, Catherine; Plossl, Karl; Skovronsky, Daniel; Gur, Tamar L.; Trojanowski, John Q.; Lee, Virginia M.-Y.; Kung, Hank F.  
 CS Department of Radiology, University of Pennsylvania, Philadelphia, PA, 19104, USA  
 SO Nuclear Medicine and Biology (2001), 28(8), 887-894  
 CODEN: NMBIEO; ISSN: 0969-8051  
 PB Elsevier Science Inc.  
 DT Journal  
 LA English  
 CC 8-9 (Radiation Biochemistry)  
 Section cross-reference(s): 28  
 AB It is well known that overprodn. and accumulation of  $\beta$ -amyloid ( $A\beta$ ) plaques in the brain is a key event in the pathogenesis of Alzheimer's disease (AD). Previously it was demonstrated that [ $^{125}I$ ]TZDM, 2-(4'-dimethylaminophenyl)-6-iodobenzothiazole, a thioflavin derivative, was an effective ligand with good in vitro and in vivo binding characteristics. To further improve the initial uptake and washout rate from the brain, important properties for in vivo imaging agents, a novel radioiodinated ligand, 2-(4'-dimethylaminophenyl)-6-iodobenzoxazole ([ $^{125}I$ ]IBOX), for detecting  $A\beta$  plaques in the brain, was **synthesized** and evaluated. The new iodinated ligand, IBOX, is based on an isosteric replacement of a sulfur atom of TZDM by an oxygen, by which the mol. weight is reduced while the lipophilicity of the iodinated ligand is increased. Partition coeffs. (P.C.) of these two ligands were 70 and 124 for TZDM and IBOX, resp. In vitro binding study indicated that the isosteric displacement yielded a new ligand with equal binding potency to  $A\beta(1-40)$  **aggregates** ( $K_i = 1.9$  and  $0.8$  **nM** for

SO Zhonghua Shenjingke Zazhi (2004), 37(6), 538-542  
 CODEN: ZSZAFN; ISSN: 1006-7876  
 PB Zhonghua Yixuehui Zazhishe  
 DT Journal  
 LA Chinese  
 CC 4 (Toxicology)  
 AB The mechanism of rotenone neurotoxicity on dopaminergic neurons was investigated. PC12 cells differentiated by nerve growth factor as dopaminergic neurons were treated by different concns. of rotenone. Cell viability was assessed with MTT, and cell apoptosis was detected by Annexin-V staining and flow cytometry. The double staining with  $\alpha$ -**synuclein** and **thioflavin T** was used to observe protein **aggregation**. After being treated with rotenone for 24 h, the process-like structures of PC12 cells disappeared, and the cell body became smaller and smoother in time- and concentration-dependent manners. Compared with the control group, the cell viability began to decline significantly when treated by rotenone at concentration of 10 nmol/L ( $A_{570} 0.415 \pm 0.013$ ) ( $P < 0.05$ ). The early sign of apoptosis was found with Annexin-V pos. staining. The apoptotic rate was  $7.35\% \pm 0.52\%$  at rotenone concentration of 5 nmol/L ( $P < 0.05$ ), and was  $13.30\% \pm 1.80\%$  at concentration of 10 nmol/L ( $P < 0.01$ ). Protein **aggregation** with the double pos. staining of  $\alpha$ -**synuclein** and **thioflavin T** were also found in the groups treated by rotenone. In vitro, rotenone should be neurotoxic to dopaminergic neurons, inducing apoptosis and inclusion of  $\alpha$ -**synuclein aggregation**. Rotenone might act through the metabolism of  $\alpha$ -**synuclein** in the pathogenesis of Parkinson's disease.  
 ST rotenone neurotoxicity dopaminergic neuron Parkinson disease

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TZDM and IBOX, resp.). Autoradiog. of postmortem brain sections of a confirmed AD patient by [125I]IBOX showed excellent labeling of plaques similar to that observed with [125I]TZDM. More importantly, in vivo biodistribution of [125I]IBOX in normal mice displayed superior peak brain uptake (2.08% at 30 min vs 1.57% at 60 min dose/brain for [125I]IBOX and [125I]TZDM, resp.). In addition, the washout from the brain was much faster for [125I]IBOX as compared to [125I]TZDM. Based on the data presented for [125I]IBOX, it is predicted that the brain trapping of this new radioiodinated ligand in the A $\beta$  containing regions will be more favorable than that of the parent compound, [125I]TZDM. Further evaluation of [125I]IBOX is warranted to confirm the A $\beta$  plaque labeling properties in vivo.

ST brain amyloid plaque imaging iodine 125 benzoxazole prepn; Alzheimer brain SPECT radioiodinated ligand prepn

IT Radiography  
(autoradiography; radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT Alzheimer's disease  
Brain  
Human  
Single-photon-emission computed tomography  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT Amyloid  
RL: BSU (Biological study, unclassified); BIOL (Biological study)  
( $\beta$ -; radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 439586-36-4P  
RL: BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 439586-38-6P  
RL: DGN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 121-88-0, 5-Nitro-2-aminophenol 619-84-1, 4-Dimethylaminobenzoic acid  
RL: RCT (Reactant); RACT (Reactant or reagent)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 118040-54-3P 439586-35-3P 439586-37-5P  
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 346691-96-1  
RL: DGN (Diagnostic use); PKT (Pharmacokinetics); BIOL (Biological study); USES (Uses)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain: comparison with [125I]TZDM)

IT 2390-54-7, **Thioflavin T** 346691-79-0 346691-94-9  
RL: BSU (Biological study, unclassified); BIOL (Biological study)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain: effect of thioflavins on [125I]TZDM binding)

RE.CNT 31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

- (1) Agdeppa, E; J Lab Compds Radiopharm 2001, V44, PS242
- (2) Agdeppa, E; J Nucl Med 2001, V42, P65P
- (3) Agdeppa, E; J Nucl Med 2001, V42, PS64P
- (4) Ashburn, T; Chem Biol 1996, V3, P351 CAPLUS
- (5) Dezutter, N; Eur J Nucl Med 1999, V26, P1392 CAPLUS
- (6) Dezutter, N; J Lab Compds Radiopharm 1999, V42, P309 CAPLUS

TZDM and IBOX, resp.). Autoradiog. of postmortem brain sections of a confirmed AD patient by [125I]IBOX showed excellent labeling of plaques similar to that observed with [125I]TZDM. More importantly, in vivo biodistribution of [125I]IBOX in normal mice displayed superior peak brain uptake (2.08% at 30 min vs 1.57% at 60 min dose/brain for [125I]IBOX and [125I]TZDM, resp.). In addition, the washout from the brain was much faster for [125I]IBOX as compared to [125I]TZDM. Based on the data presented for [125I]IBOX, it is predicted that the brain trapping of this new radioiodinated ligand in the A $\beta$  containing regions will be more favorable than that of the parent compound, [125I]TZDM. Further evaluation of [125I]IBOX is warranted to confirm the A $\beta$  plaque labeling properties in vivo.

ST brain amyloid plaque imaging iodine 125 benzoxazole prepn; Alzheimer brain SPECT radioiodinated ligand prepn

IT Radiography  
(autoradiography; radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT Alzheimer's disease  
Brain  
Human  
Single-photon-emission computed tomography  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT Amyloid  
RL: BSU (Biological study, unclassified); BIOL (Biological study)  
( $\beta$ -; radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 439586-36-4P  
RL: BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 439586-38-6P  
RL: DGN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 121-88-0, 5-Nitro-2-aminophenol 619-84-1, 4-Dimethylaminobenzoic acid  
RL: RCT (Reactant); RACT (Reactant or reagent)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 118040-54-3P 439586-35-3P 439586-37-5P  
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain)

IT 346691-96-1  
RL: DGN (Diagnostic use); PKT (Pharmacokinetics); BIOL (Biological study); USES (Uses)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain: comparison with [125I]TZDM)

IT 2390-54-7, Thioflavin T 346691-79-0 346691-94-9  
RL: BSU (Biological study, unclassified); BIOL (Biological study)  
(radioiodinated (dimethylaminophenyl)iodobenzoxazole for imaging amyloid plaques in brain: effect of thioflavins on [125I]TZDM binding)

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- (3) Agdeppa, E; J Nucl Med 2001, V42, PS64P
- (4) Ashburn, T; Chem Biol 1996, V3, P351 CAPLUS
- (5) Dezutter, N; Eur J Nucl Med 1999, V26, P1392 CAPLUS
- (6) Dezutter, N; J Lab Compds Radiopharm 1999, V42, P309 CAPLUS

- (7) Elhaddaoui, A; Biospectroscopy 1995, V1, P351 CAPLUS
- (8) Ginsberg, S; Cerebral cortex: neurodegenerative and age-related changes in structure and function of cerebral cortex 1999, P603 CAPLUS
- (9) Golde, T; Biochem Biophys Acta 2000, V1502, P172 CAPLUS
- (10) Han, G; J Am Chem Soc 1996, V118, P4506
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- (12) Klunk, W; Life Sci 1998, V63, P1807 CAPLUS
- (13) Klunk, W; Neurobiol Aging 1995, V16, P541 MEDLINE
- (14) Klunk, W; Society for Neuroscience Abstracts 1997, V23, P1638
- (15) Lee, V; Neuron 1999, V24, P507 CAPLUS
- (16) Link, C; Neurobiol Aging 2001, V22, P217 CAPLUS
- (17) Lippa, C; Neurology 2000, V54, P100 MEDLINE
- (18) Lorenzo, A; Proc Natl Acad Sci USA 1994, V91, P12243 CAPLUS
- (19) Mathis, C; J Lab Compds Radiopharm 1997, V39, PS94
- (20) Mathis, C; J Lab Compds Radiopharm 2001, V44, PS26
- (21) Mathis, C; J Nucl Med 2001, V42, P113P
- (22) Mathis, C; J Nucl Med 2001, V42, P252P
- (23) Munson, P; Anal Biochem 1980, V107, P220 CAPLUS
- (24) Selkoe, D; Alzheimer's Disease 1999, P293
- (25) Selkoe, D; JAMA 2000, V283, P1615 MEDLINE
- (26) Selkoe, D; Science 1997, V275, P630 CAPLUS
- (27) Skovronsky, D; Proc Natl Acad Sci USA 2000, V97, P7609 CAPLUS
- (28) Styren, S; J Histochem Cytochem 2000, V48, P1223 CAPLUS
- (29) Terashima, M; Synthesis 1982, V484-5
- (30) Zhen, W; J Med Chem 1999, V42, P2805 CAPLUS
- (31) Zhuang, Z; J Med Chem 2001, V44, P1905 CAPLUS

L5 ANSWER 9 OF 12 CAPLUS COPYRIGHT 2005 ACS on STN

AN 1993:444464 CAPLUS

DN 119:44464

ED Entered STN: 07 Aug 1993

TI **Thioflavin T** interaction with **synthetic**  
Alzheimer's disease  $\beta$ -amyloid peptides: Detection of amyloid  
**aggregation** in solution

AU LeVine, Harry, III

CS Dep. Neurosci. Pharmacol., Warner-Lambert Co., Ann Arbor, MI, 48106-1047,  
USA

SO Protein Science (1993), 2(3), 404-10

CODEN: PRCIEI; ISSN: 0961-8368

DT Journal

LA English

CC 9-5 (Biochemical Methods)

Section cross-reference(s): 14

AB Thioflavine T (ThT) assoc. rapidly with **aggregated** fibrils of  
the **synthetic**  $\beta$ /A4-derived peptides  $\beta$ (1-28) and  
 $\beta$ (1-40), giving rise to a new excitation (ex) (absorption) maximum at  
450 nm and enhanced emission (em) at 482 nm, as  
opposed to the 385 nm (ex) and 445 nm (em) of the free  
dye. This change is dependent on the **aggregated** state as  
monomeric or dimeric peptides do not react, and guanidine dissociation of  
**aggregates** destroys the signal. There was no effect of high salt  
concns. Binding to the  $\beta$ (1-40) is of lower affinity,  $K_d$  2  $\mu$ M,  
while it sats. with a  $K_d$  of 0.54  $\mu$ M for  $\beta$ (1-28). Insulin fibrils  
converted to a  $\beta$ -sheet conformation fluoresce intensely with ThT. A  
variety of polyhydroxy, polyanionic, or polycationic materials fail to  
interact or impede interaction with the amyloid peptides. This  
fluorometric technique should allow the kinetic elucidation of the amyloid  
fibril assembly process as well as the testing of agents that might  
modulate their assembly or disassembly.

ST thioflavine T amyloid protein fluorescence Alzheimer

IT Mental disorder

(Alzheimer's disease, pathogenesis of, amyloid fibril formation in,  
thioflavine T interaction with **synthetic**  $\beta$ -amyloid

- (7) Elhaddaoui, A; Biospectroscopy 1995, V1, P351 CAPLUS
- (8) Ginsberg, S; Cerebral cortex: neurodegenerative and age-related changes in structure and function of cerebral cortex 1999, P603 CAPLUS
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- (27) Skovronsky, D; Proc Natl Acad Sci USA 2000, V97, P7609 CAPLUS
- (28) Styren, S; J Histochem Cytochem 2000, V48, P1223 CAPLUS
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- (31) Zhuang, Z; J Med Chem 2001, V44, P1905 CAPLUS

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CODEN: PRCIEI; ISSN: 0961-8368

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variety of polyhydroxy, polyanionic, or polycationic materials fail to  
interact or impede interaction with the amyloid peptides. This  
fluorometric technique should allow the kinetic elucidation of the amyloid  
fibril assembly process as well as the testing of agents that might  
modulate their assembly or disassembly.

ST thioflavine T amyloid protein fluorescence Alzheimer

IT Mental disorder

(Alzheimer's disease, pathogenesis of, amyloid fibril formation in,  
thioflavine T interaction with **synthetic**  $\beta$ -amyloid



protein-derived peptide fragments studied by fluorometry in relation to)

- IT Proteins, specific or class  
RL: ANST (Analytical study)  
(amyloid A4, **synthetic** peptides derived from, thioflavine T interaction with, fluorometry in study of, Alzheimer's disease pathogenesis and amyloid fibril formation in relation to)
- IT 2390-54-7, Thioflavine T  
RL: ANST (Analytical study)  
(**synthetic**  $\beta$ -amyloid peptide interaction with, fluorometry in study of, Alzheimer's disease pathogenesis and amyloid fibril formation in relation to)
- L5 ANSWER 10 OF 12 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.  
on STN
- AN 86011167 EMBASE
- DN 1986011167
- TI Immunotactoid glomerulopathy.
- AU Korbert S.M.; Schwartz M.M.; Rosenberg B.F.; et al.
- CS Section of Nephrology, Department of Medicine and Pathology, Rush Medical College, Chicago, IL, United States
- SO Medicine, (1985) Vol. 64, No. 4, pp. 228-243.  
CODEN: MEDIAV
- CY United States
- DT Journal
- FS 006 Internal Medicine  
028 Urology and Nephrology  
005 General Pathology and Pathological Anatomy  
026 Immunology, Serology and Transplantation
- LA English
- ED Entered STN: 911210  
Last Updated on STN: 911210
- AB We present 11 patients with immunotactoid glomerulopathy, a new **syndrome** characterized clinically by proteinuria (11/11), microscopic hematuria (9/11) and hypertension (9/11). The patients consisted of six females and five males, aged 25 to 59 years (mean, 44.6). Proteinuria was the presenting feature and the reason for renal biopsy in all patients. The diagnosis of immunotactoid glomerulopathy was established at renal biopsy by the presence of glomerular extracellular microtubules composed of immune reactants. All the biopsies studied by immunofluorescence (10 cases) had glomerular deposits of IgG and C3. In three biopsies studied with IgG subclass specific antisera, only one patient had monoclonal immunoglobulin deposits (IgG3 kappa). In six cases the glomerular deposits were analyzed for light chains. In three the deposits contained kappa only, and three consisted of both kappa and lambda. In two cases the immune **aggregates** were confined to the mesangium, and in the remaining eight cases, the deposits were present in the mesangium and the glomerular basement membranes. Electron-dense deposits composed of microtubules were present in the same distribution within the glomerulus as the immune reactants. The microtubules had a uniform diameter in each biopsy, but they varied in size from case to case. They were approximately the same size in eight cases (mean, 22.3  $\pm$  3 [SD] nm). Three cases had much larger microtubules: 34.2 nm, 35.4 nm, and 48.9 nm in diameter. Although the 22.3-nm microtubules resembled amyloid in their appearance, glomerular distribution and random orientation in the tissue, they were more than twice the diameter of amyloid (8.9 nm), and Congo red and **thioflavin T** stains for amyloid were negative. Similar microtubular structures have been described in patients with cryoglobulinemia, SLE and paraproteinemia, but these diseases were excluded in our patients on clinical, serologic and in some case histologic grounds. More important, none of our patients had clinical or histochemical evidence of amyloidosis, an entity which may be confused

protein-derived peptide fragments studied by fluorometry in relation to)

IT Proteins, specific or class

RL: ANST (Analytical study)

(amyloid A4, **synthetic** peptides derived from, thioflavine T interaction with, fluorometry in study of, Alzheimer's disease pathogenesis and amyloid fibril formation in relation to)

IT 2390-54-7, Thioflavine T

RL: ANST (Analytical study)

(**synthetic**  $\beta$ -amyloid peptide interaction with, fluorometry in study of, Alzheimer's disease pathogenesis and amyloid fibril formation in relation to)

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on STN

AN 86011167 EMBASE

DN 1986011167

TI Immunotactoid glomerulopathy.

AU Korbert S.M.; Schwartz M.M.; Rosenberg B.F.; et al.

CS Section of Nephrology, Department of Medicine and Pathology, Rush Medical College, Chicago, IL, United States

SO Medicine, (1985) Vol. 64, No. 4, pp. 228-243.

CODEN: MEDIAV

CY United States

DT Journal

FS 006 Internal Medicine

028 Urology and Nephrology

005 General Pathology and Pathological Anatomy

026 Immunology, Serology and Transplantation

LA English

ED Entered STN: 911210

Last Updated on STN: 911210

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with immunotactoid glomerulopathy on a morphologic basis. Follow-up, from 22 to 94 months (mean, 52.6) was obtained in all 11 patients, and 2 clinical courses were noted. Six patients had progressive deterioration of renal function, with five requiring dialysis. This group had severe hypertension (4/6) and nephrotic-range proteinuria (5/6) at some point in their course. The remaining five patients with stable renal function had proteinuria of less than 2.0 g/24 hr in most cases (4/5), and none had severe hypertension. This dichotomy correlated with the distribution of immunotactoids. The patients with progressive renal insufficiency had extensive deposits involving both the mesangium and glomerular capillary walls. In contrast, the patients with less widely distributed deposits appeared to have a more stable course. Immunotactoid glomerulopathy represents a **syndrome** with characteristic morphologic and ultrastructural features. The immunotactoid microtubules are heterogeneous in size and immunoglobulin composition. Although the pathogenesis of this lesion is not known, the immunotactoids appear to represent immune reactants with a degree of ultrastructural organization which is greater than that of the various organized cryoglobulins but less than the highly structured beta-pleated sheet of amyloid. It is hoped that increased awareness of immunotactoids and further characterization of their ultrastructural composition will shed light on this newly described entity.

CT Medical Descriptors:

\*glomerulonephritis  
 \*glomerulopathy  
 hematuria  
 hypertension  
 kidney biopsy  
 proteinuria  
 cardiovascular system  
 diagnosis  
 kidney  
 priority journal  
 adult  
 etiology  
 clinical article  
 human  
 blood and hemopoietic system  
 urinary tract

Drug Descriptors:

\*complement component c3  
 \*immunoglobulin g

RN (complement component c3) 80295-41-6; (immunoglobulin g) 97794-27-9

L5 ANSWER 11 OF 12 MEDLINE on STN

AN 1998169534 MEDLINE

DN PubMed ID: 9501253

TI Alpha2-macroglobulin associates with beta-amyloid peptide and prevents fibril formation.

AU Hughes S R; Khorkova O; Goyal S; Knaeblein J; Heroux J; Riedel N G; Sahasrabudhe S

CS Biotechnology Group and the Central Nervous System Disease Group, Hoechst Marion Roussel, Inc., P.O. Box 6800, Bridgewater, NJ 08876-0800, USA.

SO Proceedings of the National Academy of Sciences of the United States of America, (1998 Mar 17) 95 (6) 3275-80.

Journal code: 7505876. ISSN: 0027-8424.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 199804

ED Entered STN: 19980422

Last Updated on STN: 19980422

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 blood and hemopoietic system  
 urinary tract

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\*complement component c3  
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Last Updated on STN: 19980422

Entered Medline: 19980410

AB We have used the yeast two-hybrid system to isolate cDNAs encoding proteins that specifically interact with the 42-aa beta-amyloid peptide (Abeta), a major constituent of senile plaques in Alzheimer's disease. The carboxy terminus of alpha2-macroglobulin (alpha2M), a proteinase inhibitor released in response to inflammatory stimuli, was identified as a strong and specific interactor of Abeta, utilizing this system. Direct evidence for this interaction was obtained by co-immunoprecipitation of alpha2M with Abeta from the yeast cell, and by formation of SDS-resistant Abeta complexes in polyacrylamide gels by using **synthetic** Abeta and purified alpha2M. The association of Abeta with alpha2M and various purified amyloid binding proteins was assessed by employing a method measuring protein-protein interactions in liquid phase. The dissociation constant by this technique for the alpha2M-Abeta association using labeled purified proteins was measured ( $K_d = 350 \text{ nM}$ ). Electron microscopy showed that a 1:8 ratio of alpha2M to Abeta prevented fibril formation in solution; the same ratio to Abeta of another acute phase protein, alpha1-antichymotrypsin, was not active in preventing fibril formation in vitro. These results were corroborated by data obtained from an in vitro **aggregation** assay employing Thioflavine T. The interaction of alpha2M with Abeta suggests new pathway(s) for the clearance of the soluble amyloid peptide.

CT \*Amyloid beta-Protein: ME, metabolism

Biotinylation

DNA, Complementary

Hela Cells

Humans

Neurofibrils

\*Peptide Fragments: ME, metabolism

Precipitin Tests

\*Protease Inhibitors: ME, metabolism

Protein Binding

Thiazoles

alpha-Macroglobulins: GE, genetics

\*alpha-Macroglobulins: ME, metabolism

RN 2390-54-7 (thioflavin T)

CN 0 (Amyloid beta-Protein); 0 (DNA, Complementary); 0 (Peptide Fragments); 0 (Protease Inhibitors); 0 (Thiazoles); 0 (alpha-Macroglobulins); 0 (amyloid beta-protein (1-40)); 0 (amyloid beta-protein (1-42))

L5 ANSWER 12 OF 12 MEDLINE on STN

AN 85239836 MEDLINE

DN PubMed ID: 4010500

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AU Korbet S M; Schwartz M M; Rosenberg B F; Sibley R K; Lewis E J

SO Medicine; analytical reviews of general medicine, neurology, psychiatry, dermatology, and pediatrics, (1985 Jul) 64 (4) 228-43.

Journal code: 2985248R. ISSN: 0025-7974.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Abridged Index Medicus Journals; Priority Journals

EM 198508

ED Entered STN: 19900320

Last Updated on STN: 19900320

Entered Medline: 19850821

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Although the 22.3-nm microtubules resembled amyloid in their appearance, glomerular distribution and random orientation in the tissue, they were more than twice the diameter of amyloid (8.9 nm), and Congo red and **thioflavin T** stains for amyloid were negative. Similar microtubular structures have been described in patients with cryoglobulinemia, SLE and paraproteinemia, but these diseases were excluded in our patients on clinical, serologic and in some cases histologic grounds. More important, none of our patients had clinical or histochemical evidence of amyloidosis, an entity which may be confused with immunotactoid glomerulopathy on a morphologic basis. Follow-up, from 22 to 94 months (mean, 52.6) was obtained in all 11 patients, and 2 clinical courses were noted. Six patients had progressive deterioration of renal function, with five requiring dialysis. This group had severe hypertension (4/6) and nephrotic-range proteinuria (5/6) at some point in their course. The remaining five patients with stable renal function had proteinuria of less than 2.0 g/24 hr in most cases (4/5), and none had severe hypertension. This dichotomy correlated with the distribution of immunotactoids. (ABSTRACT TRUNCATED AT 400 WORDS)

CT Check Tags: Female; Male  
Adult  
Amyloidosis: PA, pathology  
Basement Membrane: UL, ultrastructure  
Creatinine: BL, blood  
Cryoglobulins: AN, analysis  
Glomerular Mesangium: PA, pathology  
Glomerular Mesangium: UL, ultrastructure  
Hematuria: PA, pathology  
Humans  
\*Kidney Diseases: PA, pathology  
\*Kidney Glomerulus: PA, pathology  
Kidney Glomerulus: UL, ultrastructure  
Microscopy, Electron  
Microscopy, Fluorescence  
Microtubules: UL, ultrastructure  
Middle Aged  
Proteinuria: PA, pathology  
RN 60-27-5 (Creatinine)  
CN 0 (Cryoglobulins)

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Connection closed by remote host